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- Novel non-lonic polyurethene resins heving polyether back-bonee in water-dilutable basecoats.
- (P) This invention, therefore, relates to the field of polyurethane coatings for use in automobile basecoat/clearcoat systems. In particular, this invention relates to water-dispersible polyurethane resins having -polyether sidechains which may be utilized in basequet compositions as principal resins or grind resins. These Spolyurethane resins may be laterally stabilized or terminally stabilized. The terminally stabilized resins of this __invention are preferably employed as principal resins in basecoat compositions. 261

Novel Non-Ionic Polyurethane Resins Having Polyether Backbones in Water-Dilutable Basecoats

BACKGROUND OF THE INVENTION

Multi-layer systems have been utilized to cost automobiles for a number of years, but the early dependent of these systems necessarily employed organic solvents. As an environmental regulations became more stringent, and the cost of organic beotent rose, organic-borne systems became less desirable. The recent research emphasis in the area of multi-layer systems, especially basecost systems has tocused on the development of withshorner visitines in multi-larer confirms.

The shift from organic solvents to water for dispensing and applying resins in multi-layer systems solved many of the environmental and cost problems associated with the use of organic solvents. Water-borne systems, however, have resulted in other problems.

One solution to these problems has been the development of article polyverthere resize as described, or example, in Serial No. 003,85. These reside within speric conting characteristic may be associated accust multi-leger systems. However, in certain instances these reside may not be compatible with storage calculative of the sendance in multi-leger characteristic control of the sendancy to exhibit swinsing, and lose of ID (Districtness of Image), especially when light colored pigments (white, light blue, villes, vici.) are

The polyurchians resins of the present investion, however, may be used in the above-described motamine-outsiding systems without the negative characteristics exhibited by assionic resins. The combination of non-innic polyurchiane basecosts with storing acid catalysed clearceasts allows for a lower composite Visitate Organic Content (VICC) than with annice polyurchiane basecosts.

The retins of the present invention have excellent water deposition properties, a surprising result considering the abstence of self-forming groups. It is particularly suppring that these eaths maket an invention of the present invention of the properties where deposition years when formulated as principal retins in base cost compositions. It is also a surprising result, considering the water depositionly of these resins, that the resins of the prepart invention coalesce and hold firmly onto a ventrical panel.

In a surther aspect of the present invention the resins are formulated into grind restrict for use in basecutats. Yet, aqueous dispersions of nonhoric polyurethanee are known in the filterature and have been used to propare films and costings. Generally, polymer dispersions are unstudiable for use in milling presents. It is quite surprising that the aqueous dispersions of the nonlonic polyurethanes of the present to invention are effective ordin registers for a wide variety or diplaments.

The present invention is, in part, directed to polyure/have coatings to be used in formulating basecoat compositions of multi-layer coating systems. These reside softbit superior coating characteristics, for example, good metablic effects such as very ferentiable arrangement, sistem, and tiple factor of the metallic playments in the point film. When non-metallic playments are used, the residus of the present invention exhibit a high level of decorative effect.

This investion, therefore, relates to the field of polyurethere coalings for use in automobile basecoal/basecoa

These polyurethans coating compositions exhibit coating characteristics superior to those taught by the art and are especially useful as baseceats for automobiles.

It is an object of this invention to provide polyether-containing polyurethane regins that can be

Incorporated into baseocat formulations as principal resits and/or gried resits. It is an additional object of the invention to provide water-dispersible resine which are water-dispersible without incorporating saltas forming groups into the resins. It is an additional object of this invention to provide water-dispersible polywerbage resine which exhibit

superior coating and cosmetic characteristics.
It is a further object of this invention to provide aqueous polyurethane dispersions having favorable characteristics for formicalizing grind resins.

It is also an object of this invention to provide a method of producing the resins and basecost compositions described herein.

It is a further object of this invention to provide a method for coating a metallic or plastic substrate utilizing the resins and basecost formulations of the present invention.

These and other objects of the present invention are furthered by incorporating polyether-containing polyerethane resins into basecost formulations.

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SUMMARY OF THE INVENTION

- Two types of polyether-containing polyurethane resins are described in the present invention. We have termed these two types of resins laterally stabilized and terminally stabilized.
- s Specifically, this invention relates to a laterally stabilized polyurethane coating composition comprised
 - 1, at least one organic compound having at least two reactive hydrogen atoms:
 - 2. a nonionic stabilizar prepared by the reaction of:
- i) a monthunctional polyether with a first polyisocyenate-containing compound to produce a
 narrially canced isocyanate intermediate and
 - a compound having at least one ective amine hydrogen and at least two active hydroxyl groups;
 - 3. at least one second polyisocyanate-containing compound.
- The organic compound preferably comprises a polyester polyol, a low molecular weight diol/triol or 1s mixtures, thereof Optionally, a capping agent is employed to cap any remaining free isocyanate moleties, in addition, this investment alto relates to terminally stabilized cohumethase compositions committed of:
 - at least one organic compound having at least two reactive hydrogen atoms;
 a monofunctional polyether; and
 - a polyisocyanate.
- 20 The organic compound of the terminally stabilized polyumbare compositions preferably comprises a polyretter peptyle, a low molecular weight field endoir field, or mistures thereof. Optionally, a separate infunctional hydroxyl-containing monomer is employed for use in the polyumbare resine of the present invention. Optionally, a causing agent is employed for use in the polyumbare resine or the present invention. Optionally, a causing agent is employed to can are remaining the isocurate moistices.
- In the laterally stabilised polyumbane resins, the incorporation of the polyumbar groups function to at chain extend the polyumbane resins, in contrast, the neceptoration of the polyumbane resins are contrast, the resining the resin. The two types of polyumbane resin described above can be formulated as a value-dispress basecast composition which contrains in addition to after of the above-described resins a grief resin, a cross-finiting agent, thisotopic or healogy control agents, thickness, riginants, suffram another rise profices, benifying agent, waite, filters, surfacesors.
- agents, thickners, pigments, aluminum anctor mice particles, bestiving agents, valuer, filters, surfactants, stabilizers, plasticaters, wetting agents, dispersing agents, adhesion promotors, debamons, catalysts, and additional polymers.

 In preferred embodiments of basecoat compositions of the present invention, the terminally stabilized
 - polyurethane resin is utilized as the principal resin end the laterally or terminally stabilized polyurethane resins to utilized as the grid resin.

 After formulation, the basecust composition can be sprayed or deposited onto an automobile body, preferably, in one or two costs. Generally, the even costs of basecust are applied with a several minute fash between costs. After deposition of the basecust, before application of a thip solids, content clear cost.
 - it is generally preferred to flash about 90% of the water from the basecoat for optimum appearance and to eliminate water boil of the clearcoat.

 In both the linear and branched chain polyurethane resin, a polyecter polyo
- component.

 The polyester polyol resins described hereinabove are themselves useful on virtually any elastometic substrate, but they are particularly useful when formulated into polyorethane coatings and used in basecost formulatena for deposition onto matter or justice substrates, but people y automobile bodies.
- The polyester component may be any type, i.e., branched or unbranched, and is formed from the reaction of at least one disastecytic acid component and at least one alcohol component wherein the alcohol has at least two hydroxyl moisters.
- Virtually any carboxylic sold-containing compound having two or more carboxylic sold molettes or equivalents that are useful in synthesizing polyester compounds are useful in the present invention.
- 50 The carbonylic acid component may, of course, be comprised of short-chain disarbonylic acid compounds, long chain disarbonylic acid compounds, buring the carbonylic acid was mean compounds having at least the carbonylic acid markets and leave than 16 acrons onto in the chain. These disarbonylic acids may be skyl, alsylone, acid, alloylene, and anylene, among others. In the polyveter region of the greater investion the actionopic scid containing compound may be polyfuror-to-provide acid containing compound may be polyfuror-to-provided acid containing containing compound may be contained acid containing containi
- sa Soala with 2 or more cathony groups. A preferred carbonyle-contribining compound for use in transhed polyseter resins is fernished subplice. Bond-chair aliquit a synt (dischoppic, and compounds for reside zaide soci, solicie soci, or an equivalent alightet or aromatic and are preferred. A preferred aromatic discribonylic social is inophthalic and.

The carboyle add component may also be comprised of a tray-chall ficient-polic add component. The large-thind federate face of conting component may be an shift, siefpress, subject or similar compound, but it must be stressed that infally any isospiciate discharging and consisting compound in 200 and the stress of the compound in 200 and the conting compound in 200 and the control of conting compound in 200 and in

10 in addition to the carboyilic and containing compound, the polyseler reinin is also compressed of one or more is we relocute weight disk or to this. We have heread any compound having more han one salcohol a polyo, Polyotis may be disk jid-intended containing, finding the containing or higher sitchell-institution compounds in the case of the burnel-of-chain polyonistems. The anomal and type of this containing compounds may be varied to increase the faranching effect. A pulsared tradictoric-containing compound for the use in the burnel-of-chain polyseries is trainterlyled propose.

The polyester redin or mixture of polyester mens sized to synthesize the polyurethere resins petiently are yet only iteministed. This is effected by a yethesizing the objecter strip an except of a distribution for indicational processing the component and adortificational compound depend upon the distribution strip of the polyester compound employed. The result of this yetthers is a powerfare them to be on more the bytoticy drougs (polywateriol or polyed).

The composition of the carbonylic acid component and dyel component employed to synthesize the preferred polyester restrict is such as to provide an excess of the polycli over and above the stall number of equivarient of acid present in the minture, in other words, the researchs should be residend, and the stall-timentific proportions of the respective acid and polycli components should be adjusted to give hydroxy-emminants, polyester molecules each developed; having a hydroxyl Interfaceshylor of or most.

Monocatheorylic acid and monoalcohold: may also be used in the polyester synthesis, but these are generally utilized for the purpose of chain terminating a polyester resin. As a general rate, where used, the improcationylic telds and/or monoalcohols comprise a very small percentage by weight of the final solvester resin.

26 As a general rule the polyester dist component comprises between about 20% and 80% by weight of the final polyumrature resin. Preherably the polyester dist compress between about 50 and 70% by yeard of the polyumrature resin and most preferably the polyester dist comprises between about 55 and 56% by weight.

While it is recognised that almost any size chain length of polyester polyol can be utilized, it is preferable to use a polyester did within the melicular weight range of between 500 and 5000. It is preferable that the molecular weight range of the polyester diol component be between 1,000 and 3,500. In addition to the octivement did, the polyesteribate session of the operator to the convention are also comprised of

addition dispatic compounds having at least two relacion hydrogen storms. This compounds is primately is we recorder weight due or two compounds at lengt centres stored grounds that sold sentence or polymeric-contisting compound may be utilized at a separate compound the Thus, any did or traveloristing compound may be used. In several relationing colors of all 3 Travelorist (16) modecular contenting compound may be used. In several relationing of centre of 3 Travelorist (16) modecular contenting compounds on the settle-dispatialisty of the first polyweithers relation. The purpose of this about constraining compounds it is provided cache severalise and for fractioning compounds, provided that contention and the technique contention of the content of the co

52 The amount of low molecular weight diel andier folio stillized in the polyurethrane restine of the present invention many vary between 0 and 20 process by weight of the polyurethrane restin. Prefurably this low molecular weight alcohol component comprises between about 0 and 10 percent of the polyurethrane restin and most orderably comortises between about 1 and 6% by weight of the polyurethrane restin.

The polywerham resine of the present hvention further comprise a polyecogenee, pretensity is discognate. Generally, the discognate comprise between about 5 and 40 by weight of the final polywerham retin. Pretensity, the discognate comprises between about 10 and 30 percent by weight of the final retin and most preferably comprises between about 10 and 20 percent by weight of the polywerham retin.

A polyether-containing compound provides the polyethribate seni with the preferred wether dependability characteristics. These wither-dispenditility characteristics in usual of the polyethrane restin of the previous invention without the need to incorporate soft-forming groups within the seleri. The absence of sall-forming groups enables the polyethrane restins to be incorporated into baseasc compositions within lawy be sufficient contribution with senior action of carbonet for the polyethrane restins to be incorporated into baseasc compositions within lawy be sufficient contribution with senior part of carbonet for the provided of contribution of the provided of carbonet of the same winking, loss of I/I (distinctive lange).

"browning" effects shown by the cation-containing polyurethane regins.

The different approaches for incorporating polysither segments into the polymerhane resines are available depending upon the type of polymerhane resinised. The approach is preferred in present to previously a stabilized polymerhane utilizes a polymer of or present from the next or it is monthunctional polymer with a discontrastic to first an object with the disciplinarity of discontrast is the size of polymerhane discontrast to the size of the property of the discontrast of the discontrast of the size of the discontrast of the discontrast

10 Con symboscac, the convince installar in the macker to a maction instance comprised of a lests on organic compound having low or more scarce hydrogen functionables, and se exceeds of a phylipsoprised containing compound (in addition to that which is inceptorated into the monoise stabilizer, Preterably e polywester polyel is also added to them to polywerbane, objecting payer, for example, interestly is prepare or deshiredaminar may be used to cap say remaining their inceptorate groups. The preferably elegate or deshiredaminar may be used to cap say remaining their inceptorate groups. The preferably ellipse data a glader team for steamly stabilizer operation and containing the second or preferably ellipse data a glader team. The statings stabilizer operation are made upon the formation of the second or preferably ellipse to a glader team. The statings stabilizer operation are made to the containing the stabilization of the second or the

principal resin, but for purposes of the present invention, the terminally stabilized polyurethane resin is preferably utilized as the principal resin, and can also be used as a grind resin.

The polywher component which intalls water disputable characteristics to the laterally stabilized another characteristics to the laterally stabilized based-of-charactery polywher having not an opilwher having not uniticed promotioned by uniting a monotonic district polyment opilor. The polywher component is generally produced by utilizing a monotonic distriction of monotonic of the polywher component means the polywher component may be any group reactive with biologicalities to the national polywher component may be any group reactive with biologicalities to their a stability of the polywher component may be any group reactive with biologicalities to their a stability of the polywhere component may be any group reactive with biologicalities to their a stability of the polywhere component may be any polywhere the poly

In the laterally stabilized polywethner entit, the monthunctions playether compound it, as previously described, recision with on polywether containing companied from an laterapped locoprass. This half-capped locyclass is then reacted with a compound having an amino active hydrogen and at least to five hydroxyl compound for evaluative factorisation. The second product better did no noticel stabilizer, is bent pictoryl compound to the polywethner react by reaction with the other components. In content, in the termination stabilization polywethner resist, the polywethner resist, the polywethner inclination is uncorporated into the ends at the monthunctional stabilization polywethner resist, the polywethner resistance of the

polyether.

The manufunctional polyether compound is reacted with a mixture comprised of at least one polyester polyet, a polyticocyanate, and in addition, optionally a short chain low molecuter weight did or higher or turnional polyet, or mixtures of data and polyets. Optionally, a capping agent, for example, trimethylo-incosen or defithemed senies may be used.

For the isrminally statistical polyurethane, the polyester dol preferably comprises between 20 and 80% by weight of the final polyurethane resin and generally has a molecular weight between about 500 and 5,000, preferably between 1,000 and 3,500. Preferably, the polyester polyol component compress between

at about 50 and 70% and most preferably between about 55 and 85% by weight of the polyurethane resin. The terminally stabilized polyurethane seed in the sense weight percentages of the low wholcoals weight distribut component as the islandily stabilized polyurethane resin the Affect these abord-containing compounds are mixed, it is preferred that the ratio of tricified be higher for the terminally stabilized polyurethane resin than in the case of latently stabilized polyurethane resin than 10 and 1

This same weight pecertages of polystocyanste used in the laintally stabilized polyumbane are used in the terminally stabilized polyumbane. Preferably, the polyscoyanas is a discoyanas comprising between about 10 and 30% and most preferably between about 10 and 20% by weight of the polyumbane, in both the listerably stabilized and seminally stabilized polyumbane resins, the polyether component may be multi-incodical life and professional course beine hydroxis, tabilities. or amines with hydroxyl drough

so preferred, and preferrably it a mono or deflactional polyether with monor-functional polyethers being particularly perferred in general, water soubse polyether-doctaining compounds are useful in embodiments of the present linearition. Polyethers tormed from monacional institute opinymenzation of ethylene oxide, propolyene culdes and militaries, theself are preferred. Of course, view myint eminusty of buyllene oxide

ethers and other longer chain ethers may be incorporated into the polyether chain without adversely affecting the water dispersibility of the resins. Most preferably, ethylene oxide polymers comprise 100% of the polyether component. In general, the polyether containing component comprises between about 2 and 40% by eight of the polywethane, preferably between about 8 and 30% by weight, and most preferably s between about 10 and 25% by weight of the final polyurethane resin.

The molecular weight of the polyether-containing compounds in general ranges from about 500 to about 7000, preferably ranges from about 1000 to 4000, and most preferably ranges from about 1200 to 3000.

The polyurethanes of the present invention are advantageously storage stable and are, of course, water dispersible. The water dispersibility of the resins is controlled by the amount of polyether character to contained in the final resin particles and the hydrophobicity of the non-polyether components.

DETAILED DESCRIPTION OF THE INVENTION

- The present invention relates to a water-soluble nonionic polycrethane resin produced by incorporating a polyether-containing compound into the resin. The invention of the present application relates to lateral stabilization polycrethane resins and terminal stabilization polycrethane resins having polycther side chains. The laterally stabilized polyurethene resin is preferably comprised of the reaction product of:
- 1) a polyester polyol which is further comprised of the reaction product of a carboxylic acidzo containing product having at least two carboxylic acid functionalities and a compound having at least two alcohol functionalties:
 - 2) at least one low molecular weight compound having at least 2 alcohol functionalities;
 - 3) at least one polyisocyanate-containing compound:
- 4) a nonionic stabilizer prepared by the reaction of a monofunctional ether with a polyisocyanate as containing compound to produce a partially capped polyisocyanate followed by reaction with a compound containing at least one active arrine hydrogen and at least two active hydroxyl crouns and: 5) optionally, a capping agent is used to cap any remaining isocyanate groups that have not yet
 - reacted. The terminal stabilization polyurethane resin is preferably comprised of the reaction product of:
 - 1) a polyester polyol;
 - 2) at least one low molecular weight diol or triol;
 - 3) a polyisocyanate: 4) a triburirovy-containing manamer

 - 5) e monofunctional hydroxy-containing polyether; and 6) optionally, a capping agent,
- The polyester component is not particularly critical to the present invention, but it is necessary that the polyester component be compatible with the desired water-dependible characteristics of the revins described herein.
- Thus, the carboxylic acid component of the polyester may be comprised of long-chain dicarboxylic so acids, shortchain dicarboxylic acids, mixtures thereof or carboxylic acid equivalents such as anhydrides. factories, and polycarbonates, emong others. Long-chein monocarboxylic solds may also be used, but these are generally employed to chain terminate the polyester resin.
- The shorter chain carboxylic acid component, if used, may be comprised of a mone, di- or higher functionality carboxylic acids or a mixture of these carboxylic acids having carbon chains of 18 or tower carbon units. Monocarboxylic acids function to terminate a polyester chain and are chosen for that purpose. it is preferable that the short chain carboxyllic acid component be a dicarboxylic acid. Such preferred dicarboxylic acid compounds include, for example, adipic, szeleic, and other allphatic dicarboxylic acids. however, any dicarbitrytic acid-containing compound compatible with the goal of maximizing waterdispersibility may be utilized. Aromatic disarborytic axids may also be employed. An especially preferred to aromatic dicarboxylic acid is isophthalic scid. Alkylene and arallylene carboxylic scids may also be used. Where branched-chains in the polyester ere desired, a carboxylic acid containing three or more carboxylic acid croups, for example citric acid, may be used. A preferred parboxytic acid-containing commound of this type is trimelitic anhydride.
- The colvester resins are synthesized from the above-described carboxylic acid component and an as excess of a polyoi component. An excess of polyoi is used so that the polyester resin preferably contains terminal hydroxyl groups. The polyol compounds preferably have an average hydroxy-functionality of at least 2
 - The polytoster regin in most cases in comprised of one or more polytols, preferably a riiol. Up to about

25 pecent by weight of its pojed component may be a polycol having time or more hydrous project par molecule. Where polycis having times or more hydrous popular set observe, the result it a binarious polycene. As a general make the laterally stabilized polycenthrane receir be comprised of a polycette having no more time about 15 pecentity weight of the authoriti-constraint operquenced of a polycette having no a contral functionalise. This summed stabilization polycenthrane may be comprised of the same seather some contral polycenthrane properties of a stabilization polycenthrane may be comprised of the contral polycenthrane properties of a stabilization polycenthrane may be comprised or a stabilization polycenthrane may be comprised of a stabilization polycenthrane may be comprised on the comprised of a stabilization polycenthrane may be comprised on the comprised of a stabilization polycenthrane may be comprised on the comprised of a stabilization polycenthrane may be comprised on the comprised of a stabilization polycenthrane may be comprised to the comprised of a stabilization polycenthrane may be a stabilization polycenthrane ma

weight of the polyst component.
While it is not always deskelable to have a triol or higher multi-functional alcohol present because of the 10 tendency to form a branched chain polysester, some branching may be desirable, especially in the case where the polyseria is to be incorporated inte a branched polysertame. There may also be present and amount of microalcohol in the polyst component, particularly it larger proportions of higher functional globally set support These microalcohols serve as chall be reminders.

The dicta which are usually employed in making the polyseter resists include allaylene glycols, such as elephine glycol, propriets glycol. But have glycols and people glycols (but have glycols and as hydrogenisted bisphere) A, cyclobeaned demahavol, capolactions doil (a., the reaction product of capolactions and ethighere glycol). Ayrocaphily/indea bispherenia. And the filling have been glycold and an experimental products of vertical types and, as indicated, polysis of higher functionally may also be difficult. Such higher functional actobal care include, for example, himship-lipotrapean, retreathy-indicates, or perspectively, or when the livership logication, without philadesis and an extra contraction of the contraction of

20 as higher molecular weight polycist.
The few molecular weight dissilation are professed in the present invention are well known in the art.
They have hydroxy values of 200 or above, studyly within the range of about 1500 to 2000, such materials includes eighted ridsh, persolatively privince polycist constaining from 2 to 15 calcular plants. Examples and entire gived. 1.4-butteredol. cytodeliphatic diols such as 1.2 cytotheranedol and cytotherane diol and cytotherane diol and cytotherane diol and cytotherane diol.

projects in standy statistics polyumbara relate which as wall in bascoal composition of the present inventor, the above-decidable object in given the project of an answer of a polytocopystella, the project object of the project object is related to profitable the polyumbara related containing composed. The general synthetic approaches are officed to synthetical the polyumbara related containing composed. The general synthetic approaches are officed to synthetic the polyumbara containing composed and project synthetic approaches are officed to some of the project short of the project short object of the project short of the pro

The polyester polyol, polyisocyanete, low molecular weight dials and/or triols, and nonionic stabilizer

any by natural in the same jour, or may be rested expended, depending upon the detect restlict. Secured restored procisions select which are more crossed in stream, some they beginned and fool Secured restored processes and the secure of restored or hydroxy groups will indicease groups. Additional dates extended being at least the soften expected processes and the secure of propring groups gifted by the secure of popular countries are referred by each of the secure of the secure of the secure of the secure of popular countries of the secure of the s

program or distribunctioninis. This low molecular weight distributioning functional alcohole which are utilized as at a separate component in synthibiting the polyrevillent revision of the continued in median friend produced and produced by the continued by the continued of the continued and continued and continued and continued alcohole include timefulful programs, timefly/bitterian and portraeryfunds.

The organic polyrocognates which is reacted with the polyretar polyric and composition weight clid.

so action to limited as described is elevated by any polytocyrente, i.e., any compound containing at least two licopyrates groups, and is professibly a dislocyrente, as, hydrocation dislocyrentes or ubsettled hydrocation dislocyrentes. Many such organic discopyrates are leven in the art, including professional discopyrates, beginning of the control of the control

discoyanate (iPDI) and methylene-bite (4 cyclohaxylliscoyanate). There can also be employed iscoyanateterminated adducts of clots, such as ethylene glycol, or 1,4- butylene glycol, etc. These are formed by reacting more than one mol of a discoyanate, such as these mentioned, with one mol of a diol to ten.

longer chain disocyanate. Alternatively, the dici can be added along with the disocyanate.

While disocvanates are preferred, other multifunctional isocyanates may be utilized. Examples are 1.2.4-benzene triisocyanate and polymethylene polyphenyl isocyanate.

It is preferred to employ an aliphatic disocyanate, since it has been found that these provide better s color stability in the finished coaling. Examples include 1,6-hexamethylene disocyanate, 1,4-butylene discovanate, methylene bis (4-cyclehexyl isocyanate) and isocharone discovanate 2-tolurne

disocvariates are also preferred. Mixtures of disocvariates can also be employed. The proportions of the disocyanate, polyester,diol and/or triol component and nonionic stabilizer or

monofunctional polyeither are chosen so as to provide an isocyanate terminated intermediate polyurathane to resin. This can be accomplished by utilizing a strictionetric excess of polyisocyanate, i.e., more than one isocyanate group per nucleophilic molety (reactive with isocyanate) in the other components. The free isocyanates that remain after reaction may then be capped with a capping agent, for example, trimethylol

Langer-chain polyurethane resins may be obtained by chain extending the polyurethane chain with dol rs and/or triol-containing compounds. In addition, although it is not preferred, additional chain extending compounds having at least two active hydrogen groups for example clots, dithiols, clamines, or compounds having a mixture of hydroxyf, thiol, and amine groups, for example, alkanolamines, aminoalkyl ercaptans, and hydroxyakyl mercaptans, among others may be used. For purposes of this aspect of the invention both primary and secondary arnine groups are considered as having one active hydrogen. Alkanolamines, for

an example, otherolamine or dietherolamine, may be used as chain extenders, and most creferably, a diol in used

Examples of preferred diols which are used as polyurethane chain extenders include 1,6 haxane diol, cyclohexanedimethylol, and 1,4-butanediol. A particularly preferred diol is neopentylglycol. Polyhydroxy compounds containing at least three hydroxyl groups may also be used as chain extenders; the use of as these compounds produces branched polyurethane resins. For purposes of the present invention, if it is preferred to minimize the amount of branching in the polyurethane resin these polytrydroxy companies should be limited to a very minor component of the polyurethane producing mixture. These higher functional polyhydroxy compounds include, for example, trimethylolpropane, trimethylolethane, pentzerythritol, among other compounds.

The polyurethane resin may be chain extended in any menner using these did and tripl containing compounds or alternative compounds having at least two active hydrogen groups. Thus, these compounds may be added to the mixture of polyisocyandie, polyester, and polyether containing compound, from the stabilizer or monofunctional polyether compound), or atternatively, may react at an intermediate stage, to link two free isocyanate groups that are present at the terminal ends of an intermediate polyurethane resin.

The polyether containing compounds, as described hereinebove, are preferably mone or di-functional polyethers with mono-functional polyethers being particularly preferred. The monofunctional polyethers are preferably formed from monoalcohol initiated polymenization of ethylene oxide, propylene oxide, and mixtures thereof. A polyether compound comprised of 100% ethylene code units is especially preferred. In its most preferred embodiment, the monofunctional polyether comprises between 10 and 25% by weight of 40 the final polyurethane resin and has a molecular weight of between 1200 and 3000.

it is generally preferred that an intermediate polyurethane resin produced by reacting the polyester reain and the mixture of polyisocyanate, diolitrial mixture and polyether-containing compound be terminated with free Isocyanate groups. To accomplish this, an excess of the polyisocyanate component is used. Thus, the diolatricits, polyether component and polyester diol will all react with Isocyanate moleties to produce 45 polyurethane resins having at least some free isocyanate groups. Of course, the molar ratio of the other components will be adjusted according to the desired characteristics of the intermediate and final

polyurethane resins. in one especially desirable embodiment of the invention, a multi-functional alcohol, for example, trimethylol propage is used to terminate the reaction (cas' the tree (socyanate groups) at the desired stane se (determined by the viscosity and isocyanate groups present), thereby also contributing residual hydroxyl groups. Particularly desirable for such purposes are aminosicohols, such as othanolamine, diethanolamine and the like, since the amino groups preferentially react with the isocyanate groups present. These expoint agents are especially preferred for use in the grind resin aspect of the present invention. Multi-functional

sloohols, such as ethylene glycol, trimethylolpropane and hydroxyl-terminated polyesters, can also be ss employed in this manner. While the ratios of the components of the polyester, the multi-functional isocyanate, the diol/triol mixture, the polyether containing compounds, and the capping agent can be varied, it will be noted by those skilled in the art that the amounts should be chosen so as to avoid gellation and to produce an uncelled, unathang reaction product containing hydroxyl groups. The hydroxyl value of the final polyurethane reaction product should be at least 5 and preferably about 20 to about 200.

The amount of polyisocyanate used in the mixture is preferably between about 10% and 30% by weight of the reactants in the mixture, and most preferably between about 10 and 20%, but will vary depending s upon the polyester used and the desired molecular weight of the final polyurethane resin. The amount of polylogograpate will also vary depending upon whether it is desired to have the intermediate polygrethane terminated with free isocyanate groups or with hydroxyl groups. Thus, where it is preterred to terminate the intermediate polyurethane resin with free isocyanates for capping with trimethylopropane or diethanolamine, an excass of polylispoyanate may be used. Where the intermediate polyurethane resin is to be terminated to by hydroxyl groups, a stoichiometric deliciency of polyisocyanate may be used.

The polyurethane resins of the present invention are formulated, along with other components, into water dispersible basecost compositions which are sprayed or electrostatically deposited onto metal or plastic substrates, for example, automobile bodies. In general, a polyurethane resin, formulated as described herein, is mixed with an aminoplast resin, a polytocyanate or other cross-linking agent, a grand is regin, water, a portion of an organic solvent, pigments and a rheology control agent. Other agents may be included, for example, various fillers, surfactants, plasticizors, stabilizers, welling apents, dispersing apents, defoamers, achievion promoters and catalysts in minor amounts. In one embodiment a branched-chain

As indicated, an equecus dispersion of the polyurethane resin is utilized as the principal or major vehicle resin. In ceneral, the principal or major vehicle resin comprises between about 0 and 90% by weight of the total solids present in the basecost composition. An acceptable polyurethane resin for use as the principal resin is a resin produced from a polyester synthesized from dimer fatty acid, isophthallo acid, and 1.6 hexanedici. The resulting polyester is then reacted with a dispoyurate of isophorone, a trial and a polyether monoalcohol and a diol, for example, negpentyl glycol. The resulting polyurethane intermediate

polyester component may also be added to the basecoat composition.

as having free isocyanate groups is then reacted with trimeth ylolpropane to cap these groups. The polyurethane reaction product as described above may be mixed with an aminoniast resin or a polyisocyanate cross-linking agent. Aminoplast resins are aldehyde condensation products of melamine. urea, and similar compounds. Products obtained from the reaction of formaldehyde with melamine, urea or benzoguanamine are most common and are preferred herein. However, condensation products of other so amines and amides can also be employed, for example, aldehyde condensates of triazines, diazines, triazoles, guardicines, guaramines and alkyl and anyl substituted derivatives of such compounds, including alkyl and anyl substituted urees and alkyl and anyl substituted melamines. Some examples of such

compounds are N.N.-dimethylurea, benzourea, dicyandiamide, formoguanamine acetogusnamine, ammetre. 2-chloro-4,6-diamino-1,3,5-triazine. 6-methyl-2,4-dismino,1,3,5-triazine, 3-5-diamino-triazole, triaminopyrimidine, 2-mercapto-4.6-diaminopyrimidine, 2.4.6-triethyl triamino-1.3.5-triazine, and the like. While the aldehyde employed is most often formaldehyde, other similar condensation products can be made from other alidehydes, for example, scetaldehyde, orotonaldehyde acrolein, benzaldehyde, furfural,

and others. The amine-aidshyde condensation products may contain methylol or similar sikylol groups, and in most 40 instances at least a portion of these alicylol groups are etherified by a reaction with an alochol to provide organic solvent-soluble resins. Any morphydric slophol can be employed for this purpose, including such alcohols as methanol, ethanol, propanol, butanol, pentanol, hexanol, heptanol, as well as benzyl alcohol and aromatic alcohols, cyclic alcohols, for example, cyclohexanol, monoethers or glycols such as Cellosolves and Carbitols. (Union Carbide), and halogen-substituted or other substituted alcohols, such as 3-

45 chioropropanol. The preferred amine-aldehyde resins are etherlifed with methanol or hidshol.

Alternatively, isocrenate-containing compounds such as those previously described may be used as cross-linking agents. Generally, when polyisocyanates are used they comprise between about 1 and 50% of the basecost composition. Preferably, a fully blocked polyisocyanate is used as a cross-linking agent. In the preparation of the blocked organic polyisocyanate any suitable organic polyisocyanate may be

used. Examples include the aliphatic compounds such as trimethylene, tetramethylene, hexamethylene, 1,2propylene, 1,2-butylene, 2,3-butylene, 1,3-butylene, ethylidine and butylidene disocyanates; the cycloslintere compounds such as 1.3-cyclopentage, 1.4-cyclohexage, and 1.2-cyclohexage discoventes the aromatic compounds such as an phenylene, p-phenylene, 4.4 -cliphenyl,1,5-naphthalene and 1,4-naphss thalene dilacovanges, the allighatic-aromatic compounds such as 4,4 diphenylene methane, 2,4 or 2,6tolylene, or mixtures thereof, 4.4 soluidine, and 1,4-xylylene discovariates: the nuclear substituted aromatic

compounds such as disnisidine discovanate, 4.4-diphenylether discovanate and objectiohenylene discovanate, the triscovanates such as triphenyl methane-4,4,4 -triscovanate, 1,3,5-triscovanate benzone

and 2.4.8-trisocyanate toluene; and the tetraisocyanates such as 4.4-diphenyl-dimethyl methane 2.2, 5.5tetraisocyanate; the polymenized polyisocyanates such as tolylene disocyanate dimers and timers and the time.

Any stables disphales, cytologischel erometic sielly monosizoital and phenolic compound may be used as a blocking again an accreatance with the persent invention, such as the resumple, liver-gladinic citorolis, such as mellyst eityl, chlorently, propyl, buni, cytologisch, letpyl, cottl, cropyl 3,35-rimentyhearou, social and significant, and this like. It is exembledly all active, but as phenolication cherely depted in consortially all actives, but and the filler, the placed compounds such bod, why glocal monosities of these, other joycal monosityst with a set the filler, the placed compounds such as phenolical textual depted in settle and beautificant out on adversal glicitate founding operations. I transplan include creats, yellow, disoplanes, dishplanes, 1 things placed and 2.5-4-bobyl-1, the placed of the property of the placed of the placed of the placed and 2.5-4-bobyl-1, the placed of the placed of

Additional blocking agents include tediary hydronyl amines such as diethylwithanolamine and oximes such as diethylwithanolamine and oximes are projected from a methylority learner oximes, action coines and cyclohexanore oxime. Use of oximes and phenois is particularly useful because specific polyiscoyanates blocked with these genetic succept a makingly low immorpratures without the need for externally added unethane forming catalysts such as those described below.

The organic polyisocyanate-blocking agent adduct is formed by reacting a sufficient quantity of blocking agent with the organic polyisocyanate to insure that no free isocyanate groups are present.

A first retain may set to a used in the basecost compositions of other present investion. The grids value may be comprised of a number of water soldies provaterian result, which may be set firster or it describes to the principal or noisi vehicle investion. Set, it is particular basecost formulation a servicial contained on the province of the principal or noisi vehicle investion. As it is particular basecost formulation as servicial to the principal or noisi vehicle investion principal principal composition as particular to the principal principal composition and principal principal principal composition and principal composition

Pigment may be incorporated into the basecut composition to provide the desired consensition. This is done by misting pigments with the short-described grid resin, and in addition, accordance of the composition of the control of the composition of the control of the composition of the composition of the composition of pigment pass with the control collection control of the composition of the composition of the control of the control of the composition of the control of the con

The final pigment paste comprises about 3% to about 65% by weight of a pigment, and about 5% to about 65% by weight of a lateally terminally stabilized polyurathane restin and optionally, up to 50% by weight of a cross-liking agent.

Any standard pigment known in the six may be used with resist of the present invention so long as base appretion can be formulated without affecting the desired characteristics of the resist. Specific examples of the type staffs or prigments may be inverged or organic, for exemple, graphia, cabon basic, size chomate, sorthum chomate, busine chomate, last dynamics, flatinal dioxide, zinc oxide, combus, sorthum chomate, busine chomate, last dynamics, flatinal dioxide, zinc oxide, combus autifice, inco notice, staminum falses, zinc sutiles, philade cyseries complexes, naphthol red, quinescritores and falogeness the include prigments, among others.

Petered aluminum fields pigments are available from Silbedine Copp, Lansfort, Pennsylveria or from Cleart Werke, Cleartestahi, West Germany. The aluminum fiels pigments provide the coating with an enhanced "metalic veneer", in a preferred embodiment of the preson invention standard grade shuman subtilized with probapts as test is used. Other motalize fields pigments, for semaptic, silver may also be the metalize pigments may also be mosed with non-metalize pigments, but these are to be cavefully chosen on a not to derininh the desired medicalize diefact.

30 The national dispersions of the bissocial in competitions are dispersion in desirable what it is preferred that the decisional water here conductance reading of less than 1.3 incomment and most preferrably less than about 5 incircothers" to prevent gussing cussed by the nation of eliminar wife. Pleastered were it also crosso to outled shall that analyze our lab years. Chest evidents may also be employed with the destinated water. An exposally prelimed solvent in Ship California, which add not not to be sent of the competition and the competition of the

promote the dispersibility of individual components in the final basecoat composition (plasticizer characteristics) and for its low volatility characteristics.

A findage control agent in also preferably incorporated into the basector composition. The findage control agent can be functioned for the entiting composition and is incorporated in amounts set will be considered as meaning to the set of the composition of the composition of the composition of the composition of the control control

10 be used in embodiments of the present invention include the femile difficult compounds and the bettering depth invention made and carponarise are the hydrophical circumsposition, to result and the second compounds, for an experiment of the second compounds, and the present of the second control to the

the final basecoat composition.
In general, the particle size of the rheology control agent plays a role in the overall thisotropic as properties of these resuns. Rheology control agents in embodements of this invention are suspended in the material. It may be protoposed that the suspended rheology control agents hunching, at least in part, through

contemptic or electrostatic interactions.

In general, the particle sizes can be from less than 0.01 microns to over about 200 microns. These sizes can be adapted to develop in part the ineclopy properties sought. In appropriate diccumstances, the

ys particle state may be from about 1 to about 10 microns.
Any additional agent used, for example, surfactants, filters, wetling agents, dispersing agents, dispersing promoters, out may be incorporated into the basecoal composition. While the agents are well-wown in the prior at, the amount used must be carefully controlled to avoid adversely affecting the casing and quick-drying pharacteristics.

The final basecost composition is adjusted to a pH of between 6.0 and 8.0. Viscosity may be adjusted using delorized water. Final basecost compositions are comprised of the following components in the inclinated winth ratios:

Table I

General Description of a Silver Metallic Paint			
Ingredient	Amount (% by weight of of Solids of Final Basecoat composition)		
Polyurethane resin	20-80%		
Melamine	5-50%		
Rheology Centrol Agent 0-20% Plament (Includes Aluminum Flakes) 0-20%			
		Acid Catalyst	0-5%

e.

The basecost compositions described hereinabove can be agained to a metal or plantic substate in tone occas using the sample is an atminiser (Britis Model 60 spray gur, available from Birks so Manufacturing Corporation, (Franklin Park, Illinos), or by using other conventional spraying manus. The basecost compositions are preferably sprayed at a 50-85pst, and a reliable humidity of between 50 and 90% (potimity at 90-90% relations) humidity and a relative humidity of between 50 and 90% (potimity at 90-90% relative humidity) and a relative humidity of between 50 and

After being deposited, the basecost compositions are flash dried within a temperature range of about room temperatures to about 145 degrees F. The preferred flash temperature is about 120 degrees. The set flash conditions described herein result in about 80% of the solvents (water pixs organics) being flashed from the basecoal in this short order of time.

After the first basecost is deposited, a second basecost can be deposited over the first without drying (flash off), or alternatively, a clearcost may be deposited over the flashed basecost. It is a surprising result

of the use of the nonnotic resins of the present invention that when straight shade (non-motalic) or metallic pigments are used, one cost of basecoat might be used to provide secalism cosmetic characteristics. Any number of classroat compositions known in the art may be used. Any known ungingened or other transpresely pigmented outling agent is in principle, suitable for use as a lopocat. A typical top cost or composition creation 30-09% filter forming resin and 45-7% volicite conject colvent,

After the clear coat is created onto the basecoat layer, the multi-layer coating is then based to cross-link the polyment vehicle and to cleme be mail amount of residual writer and organic solvent from the multi-layered polyment composition. A preferred basing site involves healing the coated substrate for a profit of 10-00 minutes at a temperature of between 150 and 300 degrees P. The basing step curres the coating to a hand, durable tim.

The final multi-favor coated substrate comprises:

 a) a waterborne basecoat compositioning comprising about 20 to about 80% by weight of said basecoat composition of a polyurethane composition;

b) about 5% to about 50% by weight of a cross-linking agent,
 c) optionally, up to about 20% by weight of a rheology control agent;

 c) optionally, up to about 20% by weight of a rheology control agent dt about 5% to about 85% by weight of a pigment paste; and

a) a clear top cost. The smedien will be further described in connection with several examples which follow. These examples are shown by way of illustration of the invention and are not meant to limit the scope of the or invention. All parts and pre-notatings in the examples are by weight unless offerwhere indicated.

Example 1: Preparation of Polyester Resin A

26 A polyseter polyol reini is propored by changing a reaction vester fixes designed with a factorating column, with \$510 g (15 file) of the polyseter resin) of lespolitic accid 1920 g (15 file) filed to propore the residual column of the polyon (15 file) filed 1920 g (15 file) file

Example 2: Preparation of Polyester B

A reaction vessel is changed with 980.0 g 43.5 WGT %) of neopentyl glycol, 664.8 g (30.1 WGT %) of sophthalic acid, 84.7 g (26.5 WGT %) of adipic acid, and 50.0 g of toluene. The mixture is heated under nitrogen to 24.0° C unitio an acid uniteer of 3.5 is reached.

Example 3: Preparation of Terminally Stabilized polyurethane resin

43.0 g of methors polywhythere oxide of 2000 equipment weight, 88.3 g of IPOL set 50.0 g of glycic other PM scottes are changed to a recting and based to 15° (for 2 linear storie fringen). The inchant is of cooler to 30° c, then 252.0 g of polyeters A.5.7 g of recepting lightcut, and 25.0 g of stimethylologopase are solded. The missing is Readed to 110° cut all MOO groups there received. The interior is closed to 30° than 900 g of teopropanel is added. The missing is tentine cooler to 80° can 4200 g of decorposed water is added it never improve alignation over 10° dimites. The resulting dispersion has a solded content of the 10° can be extended on the content of the 10° can be extended on the 10° can be extended to 10° can be solded t

Exemple 4: Preparation of Terminally Stabilized polyurethane resin

44.1 g of a butanch-initized nandom cooplymer of 75% ethylone oxide and 25% propylene oxide of so do oxerzialer weight, 88.5 g of IPCI and 93.0 g of plyosi their PM acetate are changed to a reactor and hetetod is 8°C for 2 hours under nitrogen. The miniture is allowed to oxid to 30°C and 28.0 g of polysystetic A, 9.7 g of noopenly slyocal, and 25.0 g of TMP are added. The miniture is heteth to 100°C until all NOC groups have reacted. The miniture is costed to 92°C and 90.0 g of logogoant has deep in the miniture is

further cooled to 57°C and 420.0 g of delonized water is added in under vigorous agitation over 1 hour. The resulting dispersion has a solids content of 43.0% and an average particle size of 190 nm.

6 Example Sa: Preparation of Laterally Stabilized Polyurethane Resin

Linear Nonionic PE/PU "Principle Resin"

47

PREPARATION OF A METHOXYPOLYETHER DIOL

A colores del la presente del policie familia del 180 grama el methodopolishiphere giora 2000 en 1015 grama o lormativo policie. Non el meta presente interese del meta del 180 grama del 180 serio del 180 grama de

DESIDABATION OF A POLYCIDETHANE RESIN

A flask is charged with 800 game of polyeeth A (seample * 1), 100 game of cabrowax did; 90.2 game of responser by glores; 240 game of inceptioned abcoquestis and 250 game of propriets glores; 240 game of propriets and propriets game of ga

Example 58 GRIND RESIN - LATERALLY STABILIZED NONIONIC PEPV

PREPARATION OF A CARROWAY DIOL

A cabone doll is prepared by charging a fast with 0.038 game of methopolywhyther glycal 2000 and 1015 game of blame. The midnes is behald to make to more the water present in the cabone. When did not be settle present in the cabone. When did not settle present is the prepared to \$0° care 4.5 games of \$0° ca

PREPARATION OF A POLYURETHANE RESIN

55 A flask is chapped with 800 grams of polyseler A (example = 1), 100 grams of carbonax dol, 50.2 grans of caspenty) glyco. 24-02 grams of complorume discopurate, and 205 grams of prophysical property of the carbonax of prophysical property of the carbonax dol, 50 grams of prophysical property of the carbonax dol, 50 grams of carbonax dollars in a coloration story and the carbonax dollars. Then the business of property of property of property dollars in a coloration story of the carbonax dollars.

delonized water is added under high agitation to form a dispersion.

Example 6: Preparation of a Terminally stabilized Polyurethane Resin

BIS.5 of polyseare A. 2010 of methanol-initiated polyethylene oxide of 1260 equivalent weight, 1455 of pl RV and related to 126° contained to 126° contained

Example 7: Preparation of Terminally Stebilized polyurethane resin

455.0 g of polyester B, 155.0 g of methand-initiated polyethylene oxide of 1450 equivalent weight, 10.0 g of TMP, 131.0 g of TMPC, and 132.0 g of MPK are charged to a reactor and heated to 16°C. Love introgen for six hours, at which them on NCO groups remain unnecent. 2000 g of monotouthy (glycol ether is added and the mixture is coded to 70°C. Nact, 1370.0 g of decinited water is added in over 10 mixture with respect to 10°C. Nact, 1370.0 g of decinited water is added in over 10 mixture with respect to 10°C. Nact, 1370.0 g of decinited water is added in over 10 mixture.

Example 8: Preparation of silver metallic basecost using laterally stabilized nonionic resin

Component	Parts by Weight	WGT %.NV
2% Laponito paste	242.2	3.2
Resin from example 5	201.1	47.7
Aluminum paste (ALCOA 87575)	32.9	14.9
Phosphate eater solution	1.4	0.9
Melamine (Cymel 303)	\$1.1	44.5
p-Toluenegulfonic acid catalyst (amine blocked)	9.1	1.5
deicrized water	162.2	

The polyurchane resin from example 5 is slowly added to the 2% Lipponite paste under vigorous agitation, in a separate container the aluminum stury is prepared by miding the aluminum paste, phosphase ester solution, and melamine under agitation. The similarm stury is slowly added to the resist mixture under high agitation. Delicrated water is added to medute the viscosity of the paint to 14 seconds (#2 SRSHE CUPT. No eld of the earlier is 5.7.

Example 9: Preparation of silver metallic basecoat using terminal stabilization nonionic resm

The paint is prepared according to the process of example 8 except the terminally stabilized colvurations resin of example 4 is used. The paint has a viscosity of 14 seconds (62 FISHER CUP).

Comparison study: Lateral stabilization vs. terminal stabilization silver metallic basecosts

The silver metallic basecoats from examples 8 and 9 were sprayed dich-by-side using a sighton activated spray gan bits parties was respond at 65 PSI over primed steel parels at 52 P and 46% installed humidity. The panels were based at 250 degrees F (or 50 minutes and the metallic effect was revaluated by comparison with a series of five standard silver metallic panels (1 = bost). Restults were as this results were as the parel of the panels of th

EP 0 354 261 A1

Silver Paint	Metallic Effect (1 = best)
example 8	3.0
example 9	5.0

The silver metallic basecosts from examples 6 and 9 were resprayed at 86 degrees F and 50% relative humidity to safe for the sage relatance of the partie. After basing the protect for 50 minutes at 250 degrees, a relative sag resistance value was assigned to each panel (1 = best; no sag, 5 = worst, excessive sag). Results are as follows:

Silver Paint	Sag Resistance (1 = best)
example 8	1
evennie 9	- 5

The following examples of pigment grind pastes and straight-shade paint are applicable to both types of urethanes (i.e., lateral stabilization and terminal stabilization).

Example 11: Preparation of a TiO₂ pigment paste

Dispersion of TiO₂

The nonionic prothers dispersions have been used to mill TiO₂ for example, as supplied by DuPont, Glidden, etc., and more specifically, using DuPont R960HGHG as in the example below:

	Grams	Grams Non-Volatile
Nonionic Urethane Dispersion	1452	450
DuPont R960HGHG	2250	2250
Deionized Water	331	

The unshare dispersion is placed in a two gallon vessel accipron with a propellor type agilator. The YTO is added to the stirred deposition. Debracke where is used to maintain a hald paster. After the pigment is added, the skery is stirred for 50 minutes. The viscosity is equalled to 75-68 foots units (700-1000 ccs) with words. Sitting is continued for 15 minutes. The posses is left through paywhy for sendant activation from Chicago Bolter Company, charged with ceramic or glass media. Particle states of 0-5.5 microses is readered in one or two cases.

Example 12: Preparation of a Blue Pigment Paste Dispersion of Phthalocyanine Blue Pigments

The nonionic unsthane dispersions have been used to mill phthalocyanine pigments for example, shose supplied by Ciba Golgy, Toyo Ink, Sun Chemical, etc., and more specifically Ciba Golgy XX485 as in the example below:

	Grams	Grams Non-Volatile
Nonionic urethane dispersion	1781	552
Cha Geiny X3485	276	278

The urethane dispersion is placed in a two gallon vessel equipped with a propellor type agistant. The dry pigment is added to the stimed dispersion. After the addition of the pigment, the sturry is stimed 30 minuses. Dolinion's value may be added in excessary to devise a viscosity of 60-85 kness units (500-1000 cps). Stiming is then confirmed for 15 minutes. The passe is then fad through a gravity fed eardmill as described above unit the cardict size is 0-12 minors.

Example 13: Preparation of a Perylane Figment Paste Dispersion of Perylane Figments

The nonionic urethane dispersions have been used to mill perylene pigments as supplied by Mobay, 8ASF, etc., more specifically Mobay R8424 in the example below:

	Grams	Grams Non-Volatile
Nonionic urethane dispersion	2419	774
Mobay R6424	581	581
Delicnized Water	109	581

The urethrane dispersion is charged into a two gellon vessel equipped with a propellor type egitator. The od by pigment is added to the stimed dispersion. Stiming is continued for 30 minutes after the pigment has been added. Delonitated water is used to obtain a viscosity of 50-55 Krebs units (200-5000) cps). The starry is then added to an attribut as supplied by Union Process, Alexon, Ohio, charged with stainless steel shot. After 4-8 hours the particle size is 615 micross.

Example 14: Preparation of a Carbon Black Pigment Paste Dispersion of Carbon Black Pigments

The nonionic urethane dispersions have been used to mill carbon black pigments from De Gussa, Columbian Chemicals, Cabot, etc., more specifically Cabot Black Pearls 1300 in the example below:

	Grams	Grams Non-Votatile
Nonionic Urethane Dispersion	2800	896
Cabot Black Pearls 1300	224	224

The unothane dispersion is charged with a two gallon vessel equipped with a propellor type agistor. Similarly programs is added to the dispersion. Similar is continued for 50 minutes after the pigment has been added. If necessary delocated waters is used to boths a visicosity of 50-56 fines bent 500-500 cpp.) The sturry is then added to an attribor as supplied by Union Process, Alaron, Ohio, charged with stainless steel shot, After 4-6 hours the pacific size is 6-12 minoras.

55 Example 15

		# Pigment	# Vehicle	100#
A)	Laponite Pitriol P900 DI H 0 Total for A	2	2	13.9
В.	Melamine (Cymel 303) Butyl Cellosolive		31.4	6.53 1.62
C.	Nonionic urethane Dispersion	1	33.5	21.7
D.	Furned Silica (R-872 from DeGussa) Melamine Nonsonic dispersion Total	9.5	4.61 8.74	16.2
E.	White grind paste (Example 11) Oxazolidine blocked pTSA	100.0	19.8	38.2

- Part A is mixed under high agitation for 2 hours. Part B is added slowly under egitation followed by Part
- Part D is ground separately in a mill and it and parts E and F are added under agitation. Final adjustments in viscosity are made with delonized water. The pH of the basecost is 6.0-a.D.
- The invention has been described in detail with particular reference to preferred embodiments thereof, but it will be understood that variations and modifications can be affected within the spirit and scope of the invention and that the scope of the invention is to be determined by the claims accorded hereis.

Claims

- A laterally stabilized polyurethane resin comprising the reaction product of:
- a) at least one organic compound having two or more reactive hydrogen functionalities;
 b) a nonionic stabilizer prepared by the reaction of:
- 35 i) a polyether having at least one scrive hydrogen functionality with a first polyisocyanate compound to produce a partially capped isocyanate intermediate; and
 - ii) a company deprine indexpansion in institution and at least two active hydroxyl groups; and cl at least one sective sinine hydrogen and at least two active hydroxyl groups; and cl at least one second isocvanists—containing compound.
- The polyurehane resin according to claim I wherein said organic compound is selected from the group consisting of polyester polyols, dipls, tricis and mixtures, thereot.
- The resin according to claim 1 further reacted with a capping agent.
 The resin according to cleim 1 wherein said organic compound a) is selected from the group consisting of diols, bride and mixtures thereof.
- The resin according to claim 2 wherein said organic compound a) is selected from the group consisting of ethylene glycol, propylene glycol, butylene glycol, neopentyl glycol, and 1,8-hexanediol.
- The resin according to claim 4 wherein said alcohol-containing compound comprises at most about 20 percent by weight of said polyurathane resin.
- The resin according to claim 1 wherein said second isocyanate-containing compound comprises about 10% to about 30% by weight of the final polyurethane resin.
- 8. The resin according to claim 1 wherein said polyether has a molecular weight between 1200 and 3000.
- 9. A terminally stabilized polyurethane resin comprising the reaction product of:
 - a) at least one greanic compound having at least two reactive hydrogen functional ries:
 - b) a polyether containing at least one active hydrogen; and
 - c) a polyisocyanate.
 10. The polyurethane resin according to claim 9 wherein said polyurethane resin is additionally.
- ine polyuretnane resin according to claim e wherein said polyuretnane resin is accruonal comprised of at least one trifunctional hydroxyl-containing monomer.
 - 11. The polyurethane resin according to claim 9 wherein said organic compound is selected from the

- group consisting of polyester polyols, dials, trials, and mixtures, thereof.
 - The resin according to claim 11 further reacted with 8 capping agent.
- 13. The resin according to claim 9 wherein said alcohol-containing compound comprises at most about 20 percent by weight of said polyurethane resin.
- 14. The resin according to claim 9 wherein said polylsocyanate comprises about 10% to about 30% by weight of the final polycrethane resin.
- 15. A basecost composition for spraying or depositing onto metal or plastic substrates comprising: a) a laterally stabilized polyurethane resin comprising the reaction product of:
- i) at least one organic compound having two or more reactive hydrogen functionalities;
 ii) a nonionic stabilizer prepared by the reaction of:
- i) a polyether having at least one active hydrogen group with a first polyisocyanate compound to produce a partially capped isocyanate intermediate; and
- persany cappes socyaniae intermeticas; and 2) a compound having at least one active amine hydrogen and at least two active hydroxyl groups; and iii) at least one second colvisocyanate compound:
 - b) a cross-linking agent,
 - c) a grind resin; and
 - d) a pigment component.
- 16. The polyurathane resin according to claim 15 wherein said organic compound is selected from the group consisting of polyester polyols, diols, triols and mixtures, thereof.
- 20 17. The basecat composition according to claim 15 further comprising a rheology control agent.
 18. The basecat composition according to claim 16 wherein said polyether has a molecular weight of
- 18. The basecost composition according to claim 16 wherein said polyether has a molecular weight of between 1200 and 3000.
 19. The basecost composition according to claim 15 wherein said polyurethane resin is further reacted.
- with a capping agent.
 25 The basecoat composition according to claim 15 wherein said organic compound is selected from
 - the group consisting of polyester polyols, diols, triols and mixtures thereof.

 21. The basecual composition according to claim 20 wherein said alcohol-containing compound comprises at most about 20 percent by weight of said polywerhane rests.
- The basecost composition according to claim 15 wherein said second isocyanate-containing so compound comprises about 10% to about 30% by weight of the final polyurethane resin.
 - 23. A baseouat composition for spraying or depositing onto metal or plastic substrates comprising: a) a terminally stabilized polyurethane comprising the reaction product of:
 - i) at least one organic compound having at least two active hydrogen groups;
 ii) at least one polyether having at least one active hydrogen functionality;
- as lif) a polyisocyanatic
 - b) a cross-linking agent
 - c) a grind resin; and
 - d) a pigment component
 24. The composition eccording to claim 23 further comprising at least one triflunctional hydroxyl-
- 25. The basecost composition according to claim 24 wherein said polyether has a molecular weight of between 1200 and 3000.
- 28. The basecost composition according to claim 23 wherein said polyurethane resin is further reacted 49 with a capping agent. 27. The basecost composition according to claim 23 wherein said alcohol-containing compound
- 27. The basecost composition according to claim 23 wherein said alcohol-containing compound comprises at most about 20 percent by weight of said polyurethene ratio.
 28. The basecost composition according to claim 23 wherein said grind resin comprises a laterally
- stabifized polyurethane resin according to claim 1.

 20. The baseoust composition according to claim 23 wherein said grind resin comprises a terminally stabifized polyurethane resin according to claim 9.
 - 30. A multi-layer composition comprising:
 - A waterbome basecoat composition comprising:
 The laterally stabilized polyurethane resin from claim 1;
- ss ii) a cross-linking agent; itil a pigment and
 - b) A clear toposat composition overcoating said basecoat composition.

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- 31. The multi-layer compositon according to claim 30 further comprising a rheology control agent.
- 32. The multi-layer composition according to claim 30 wherein said basecoat composition further comprises a grind resin according to the polyurethane resis of claim 1.
- The multi-layer composition according to claim 30 wherein said basecoat composition further comprises a grind retin according to claim 15.
 A multi-layer composition composition
 - a) A waterborne basecoat composition comprising:
- i) the polyurethane resin from claim 17:
- to ii) a cross-linking agent

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- ili) a rheology control agent, and
- iv) a pigment and
- b) A clear topcoat composition overcosting said basecoat composition.
- 35. A proment paste for use in waterborne basecoat compositions comprising:
- a) about 3% to about 65% by weight of said pigment paste of a pigment,
- b) about 5% to about 35% by weight of said pigment pasts of a branched chain polycrethane resin produced by the reaction of:
- i) at least one polyester polyol;
- 39 (i) at least one low molecular weight alcohol-containing compound having at least two alcohol functionalities; iii) at least one tifunctional hydroxyl-containing monomer;
 - ii) at loast one transcrional hydroxyr-containing flond
 iv) a polyether alcohol; and
 - v) a polylsocyanate; and
 - c) about 0% to about 50% by weight of said pigment paste of a cross-linking agent.
 - A multi-coated metal or plastic substrate comprising a substrate coated with a) at least one waterborne basecoat composition comprising:
 - about 20 to about 60% by weight based on the final solids contant of said basecoat composition of a linear polygrafiane composition produced by reacting:
- 1) at least one polyester polyei;
 2) at least one low molecular weight alcohol-containing compound having at least two alcohol functionalities;
 - a nonionic stabilizer prepared by the reaction of:
 a a monofunctional ether with a first isocyanate-containing compound to produce a partially capped
- isocyanate intermediate; and b. a compound having at least one active amine hydrogen and at least two active hydroxyl groups; and 24 at least one second isocyanate-containing compound.
- ii) about 5% to about 50% by weight of a cross-linking agent selected from the group consisting of aminoplass restin or polytocoparate;
 iii) about 7% to about 25% of a resolvery control asset selected from the croup consisting of timest visitors.
- components, bentonite clays, and hectorite clays;

 (iv) is about 5% to about 85% of a pigment paste; said basecost composition being flash-dried before being coated with
 - b) A clear topcoating, said basecoat and said topcoat being cured to a hard, durable film.

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EUROPEAN SEARCH REPORT

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